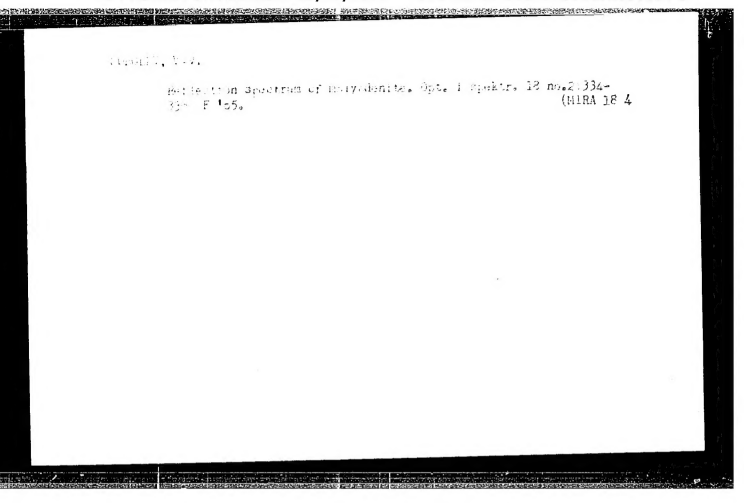


P1-4 IJP(c) ENT(1.)/ENT(m)/ENT(m)/T/ENP(t)/EEC(b)-2/EMP(b) L 49035-65 RDW/JD/GG S/0181/65/007/003/0914/0915 AP5006905 ACCESSION HR: AUTHOR: Soboler. V. V. TITIE: Line spectra of edge absorption and emission of cadmium telluride SOURCE: Fizika tverdogo tela, v. 7, no. 3, 1965, 914-915 TOPIC TAGS: cadmium telluride, group II element, group VI element, line spectrum, absorption edge, emission edge ABSTRACT: In view of the little attention paid to the edge absorption and emission of crystals of the AIBVI group, other than CdS and CdSe, the authors investigated the low-temperature absorption of thin layers and photoluminescence of singlecrystal and polycrystalline CdTe in the region of the intrinsic absorption edge. At 77.3K, the absorption spectrum of CdTe films consisted of several broad lines at 7685, 7757, and 7817 A, with a noticeable step on the continuous edge absorption curve at '7830 A. A reduction of the temperature to 4.2K shifts the entire spectrum by ~ 37 Å (0.008 eV) to the short-wave region, and reduces the line width At 4.2K the single crystal's emission spectrum consists of three narrow lines and

49035-65 ACCESSION NR: AP50069				
ly). In the case of 18 (7660, 7750, 7855, and 8330 A). An analysis of those of CdS and CdSe to free exciton, excitchas: 1 figure.	o, 8055, 8120 and 8385, 8520 arge-crystal layers the spe 7915 Å) and three bands (8 of the results shows that to than assumed by other workers, or exciton-impurity characters.	000 and 8085 [doub he spectra are much rs, and exhibit sin racter of the line	Let], 8190, and n closer to milar relation s. Orig. art.	
ASSOCIATION: Institut	fiziki i matematiki AN MSS	R, Kishinev (Insti	tute of Physics	
and Mathematics, AN MS	SR)			-2-2
and Mathematics, AN MS	ERCL: 00	SUB CODE:		
and Mathematics, AN MEN	3K)	SUB CODE:		
and Mathematics, AN MASS	ERCL: 00	SUB CODE:		
and Mathematics, AN MASS	ERCL: 00	SUB CODE:		



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61674-65 EWT(1)/EWT(m)/T/EWP(t)	/EWP(b)/EWA(c) Pi-4 IJP(c) JD
CCESSION NR: AP5011119	535.373.1
UTHOR: Sobolev, V. V.	UR/0051/65/018/004/0648/0655 535.373.1
	Me stre colonide and zinc telluride
	e of zinc selenide and zinc telluride
OURCE: Optika i spektroskopiya, v	r. 18, no. 4, 1965, 648-655
	scence, edge photoluminescence, exciton state,
OPIC TAGS: zinc compound, immines	acture, line structure, crystal structure
	tra of the edge photoluminescence of ZnSe and gere found to have complicated line and band
Man James Home Twenty	her eithlimation and the photologicalce was in the
	in Aloist and fifthatoter litters. The proofer
	th a large-aperture spectrograph. The measure- Three types of ZnSe radiation and two types of
An in the transport of	anolvais or the intensity and mail-winds are an analysis
the house and the house	indicates that edge luminescence was observed. e of the layers is deduced from a comparison of
he nature of the crystal structure he obtained spectra and earlier de	ata. It is concluded that exciton lines are more
He obtained speaker	
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7명 보다가 한다다는 생활하다면 그리다면서 다른생활하다	

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effectively produced in layer, and 5 tables.	s than in sing	e crystele,	Grig. art. b	us: 2 figu	res
ASSOCIATION: None					
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L 64505-65 EPA(s)-2/ ACCESSION NR: AI	EVA(c)/EVT(1)/EVT(m)/EVG P5012607 JG	(m)/FWP(t)/T/EWP(t)IJP(c) UR/0051/65/018/005/08 535.512:535.33 + 535.3	A The second of
AUTHOR: Sobolev		2141/5	45 B
TITLE: Reflection	on and absorption spe	ectra of crystals of the	ne
Group A ^{II} B ^{VI}			
SOURCE: Optika	i spektroskopiya, v.	18, no. 5, 1965, 813-6	319
TOPIC TAGS: cadr duction band, abs spin orbit coupl:	sorption spectrum, 1	ry \mathcal{H} compound, valence by Ight reflection, absorp	and, con- otion edge,
agonal (wurtzite spectra of hexago (sphalerite) poly single-crystal Ho) CdS and cubic (sphonal (wurtzite) sing yerystals of HgS (cy gTe and CdTe were in	polycrystalline layers alerite) CdTe, and the le-crystal CdS and CdS nnabarite) and HgSe and vestigated at 290 K and uced by sputtering of	reflection and cubic i of i 1 6 eV
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A C	at 10 mm Hg. The surfaces, of surfaces, cleaved surfaces, cleaved surfaces, cleaved surfaces, controlled the surfaces, co	at 10 ⁻⁵ mm Hg and on glass substrates were measured for reflection from natural or polished surfaces. The peaks of the or polished surfaces. The peaks of the ed and their energies calculated and tab- ed for different orientations. The ed for different orientations. The effection of the crystals are interpreted if e observed spectra. The distance cal- i and the lower conduction band cal- i and the lower conduction band. Orig. art. has: 3 figures, 1 formula, ENCL: 00 SUB CODE: OF	
	Card 2/2	\(\frac{1}{2}\).	-

RDW/JD EWT(m)/ETC/EWG(m)/EWP(t)/EWP(b) IJP(c) UR/0051/65/019/001/0086/0096 I. 11115-66 ACCESSION NR: AP5017896 535.37:548.0 44,55 AUTHOR: Sobolev, V. V.

Edge luminescence of of CdS, CdSe, and CdTe crystals TITLE:

1965, 86-96 v. 19, no. 1, Optika i spektroskopiya, SOURCE:

cadmium selenide, cadmium sulfide, cadmium telluride, TOPIC TAGS: luminescence spectrum, excitron, impurity center

ABSTRACT: In view of the contradictory character of the numerous investigations on edge luminescence, which were limited for the most part to CdS, the author attempted to detect and investigate edge luminescence spectra of other crystals isomorphous with CdS, namely CdSe and CdTe polycrystals. Pure coarsely-crystalline layers were prepared by vacuum sublimation on heated quartz substrates. Fluorescence was excited by an SVDSh-1000 lamp and photographed by reflection There was no luminescence at in spectrographs in the 700 A region. room temperature, but bright edge photoluminescence appeared at 77.3K,

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ACCESSION NR: AP5017896

and even more at 4.2K. The changes in the line and band spectrum occurring upon cooling each of the crystals are described. Based on the character of the intensity distribution, the half-width of the lines and bands, and the wavelength distribution, the photoluminescence spectra can be distinctly divided into three groups: (1) emission lines that coincide in resonance with edge absorption lines; (2) the remaining emission lines on the longer-wavelength side; (3) the edge-band emission. The lines of the first group in the edge luminescence of the crystal can be attributed to resonance radiative decay of excitons with a shift of the annihilation lines towards longer wavelengths. The second group is due to various satellites of the exciton and bound exciton-impurity states. Arguments in favor of a close connection between excitons and the tand edge luminescence are presented. The author tnanks R. Yu. Khansevarov, L. P. Bogomazor, and N. Lazovskaya for much assistance in preparing the polycrystalline layers. Orig. art. has: 3 figures and 5 tables.

ASSOCIATION: None

SUBMITTED: 02Apr64

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ENCL: 00

SUB CODE: OP, 55

NR REF SOV: 017

OTHER: 009

Card 2/2 /

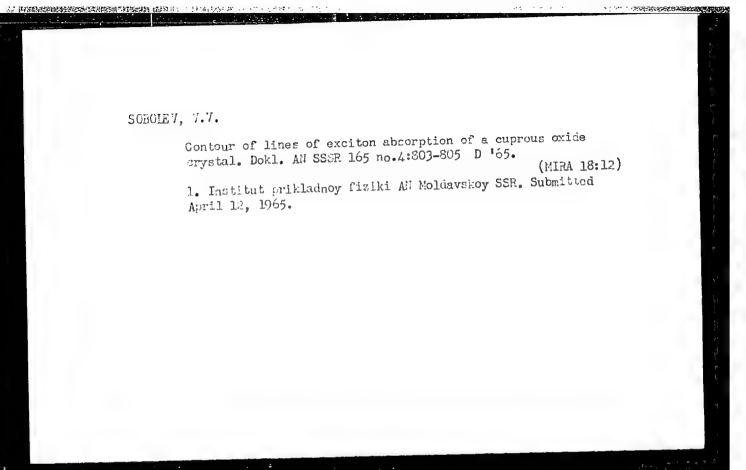
CIA-RDP86-00513R001651830004-3"

IJP(c) L 2373-66 EMT(1)/T UR/0020/65/163/004/0868/0869 ACCESSION HR: AP5020827 AUTHORS: Kesamanly, F. P.; Kroitoru, S. G.; Rud', Yu. V.; Sobolev, V. V.; Syrbu, 419.00 N. N. TITLE: The energy band structure in crystals of the group A B C 2 SOURCE: AN SSSR. Doklady, v. 163, no. 4, 1965, 868-869 TOPIC TAGS: semiconductor, zinc compound, conduction band, Brillouin zone ABSTRACT: Investigations were made of the energy structure in minerals having the structure of chalcopyrite. The lowest conduction band is simple, and the highest valence band is triple. This paper examines the reflection spectra of ZnSnAs2, ZnSiP2, and ZnSiAs2 in the region of 1-6 ev and at 293K. The spectral distribution of reflectivity showed two interes maximums for each crystal: at 265 and 600 mm for the first, 280 and 330 mm for the second, and 275-295 and 370 mm for the third. The peak at 600 mm for ZnSnAs, has a doublet structure with two maximums at 550 and 650 mm. Spin orbit splitting for ZnSnAs2 proved to be 5-10 times that for the other two. Because of the width of the peaks, doublet structure of a long-wave maximum was not observed in the reflectivity curves of the last two crystals. In Card 1/2

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L 2373-66 ACCESSION NR: AP5020827	9
general characteristics, the three minerals are very similar the reflectivity spectra are due to allowed cross-over interpoints in the Brillouin zone analogous to points L and X in	rzonal transitions at crystals of group
A ^{iv} and A ⁱⁱⁱ B ^v . The great general and detailed similarity of the tested crystals to the groups A ^{iv} and A ⁱⁱⁱ B ^v strongly	in reflectivity spectra y suggests a great
similarity in structure of the energy bands and the nature both groups. "The authors express their thanks to Professor support of the present work." Orig. art. has: 2 figures and	of the chemical bonds of r D. N. Nasledov for his
ASSOCIATION: Fiziko-tekhnicheskiy institut im. A. F. Ioffe (Physical and Technical Institute, Academy of Sciences SSSR fiziki, Akademii nauk MSSR (Institute of Applied Physics, A); Institut prikladnoy
SUBMITTED: 15Jan65 ENCL: 00	SUB CODE: SS
NO REF SOV: 005 OTHER: 003	
8 V K Card 2/2	



ACC NRI AP502890	T(1)/EWT(m)/T/EWP(t)/EWP(b)/EWA(c) IJPO 9 SOURCE CODE: UR/0020	
AUTHOR: Sobole	V, V. V.	54
(Institut prik)	of Applied Physics, Academy of Sednoy fiziki Akademii nauk MSSR)	ciences MSSR
TITLE: Low ten	perature spectrophotome in investe edge of intrinsic absorption of crystals	tigations in the region cadmium selenide and
SOURCE: AN SSS	R. Doklady, v. 165, no. 3, 1965,	534-536
TOPIC TAGS: ab spectrophotomet strength, excit	sorption edge, cadmium selenide, cric analysis, absorption coefficient	carmium sulfide, E.t. Pscillator
ABSTRACT: The coefficient of	author investigated the spectral cabsorption of pure and highly per	distribution of the fect thin (d = 0.2
	ystal hexagonal plates of CdSe (A	
on the contours	4790 Å) for the purpose of obtains of the exciton absorption bands. t at 4.2K and 160K in polarized 1	The investigations
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T. 13124-66

ACC NR: AP5028909

(D \approx 3.2 Å/mm in second order) and prism (D \approx 50 Å/mm in the region λ 5000 Å) spectrometers in special cryostats described by the author earlier (Izv. AN MSSR No. 11, 1963). The receiver was a photomultipler (FEU-17) cooled with liquid nitrogen. In the case of CdSe, the principal terms of three exciton lines could be separated at 6960, 6878, The absorption coefficients for these lines are measured and 5600 A. and the half widths and oscillator strengths calculated and tabulated. The presence of three nonoverlapping exciton series in CdSe is confirmed. Indirect transitions are observed in a small region of frequencies near the edges of the direct transitions. From a comparison with theory it is concluded that most lines of the line edge absorption of CdSe actually should have an exciton origin. The line contour points to a strong exciton-phonon interaction at 160K and a weak interaction at 4.2K. Similar measurements were made for CdS by plotting the transmission curves of different pure crystals, and it is deduced that exciton-photon interaction plays an important role in CdS at 4.2K, in contradiction to the experimental and theoretical results by others. This report was presented by Academician V. P. Konstantinov. art. hast 3 figures and 1 table.

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	TIP(c) ID	
	L 14852-66 EWT(m)/EWP(t)/EWP(b) IJP(c) JD ACC NR: AP6001725 SOURCE CODE: UR/0020/65/165/004/0803/0805	1
-	AUTHORS: Sobolev, V. V.	
	ORG: Institute of Applied Physics, Academy of Sciences MSSR (Institut prikladnoy fiziki Akademii nauk MSSR)	
	TITLE: Contour of the exciton absorption line of a cuprous oxide crystal	
	SOURCE: AN SSSR. Doklady, v. 165, no. 4, 1965, 803-805	
	TOPIC TAGS: exciton absorption, cuprous oxide, absorption line,	
	ABSTRACT: The author points out that although Cu ₂ O has been thorough	
	ly investigated in many respects, there are not enough that width, osfundamental quantities as the absorption coefficient, half width, osfundamental quantities as the absorption of its spectrum. The author cillator strength, and the line contour of its spectrum.	
	persion 3.2 A/mm, at temperatures of 160 and 4.2K, the spectral dis-	
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14852-66 ACC NR: AP6001725

tribution of the absorption coefficient of single-crystal and polycrystalline $Cu_{2}0$ with thickness 5 -- 300 μ . The exciton lines of the *yellow* series with quantum numbers n = 1 - 6 and n = 1 - 7 were observed for 160K and 4.2K respectively. The green series of lines with n=2 and 3 at 160K and n=2, 3, 4 at 4.2K were also observed. An examination of the line contours shows that all the green lines and many of the yellow lines are strongly asymmetrical, and that the asymmetry increases with n. This indicates that the interaction between the excitons and the phonons changes from weak to strong with increasing n. When the temperature is lowered from 160 to 4.2K, the absorption intensity varies in a highly irregular fashion for the different lines of the green series. The absorption at the maximum increases strongly for n > 3 and changes little for n = 2 and 3. The results are compared with those by others and many differences are attributed to differences in the spectrometer dispersion and in the quality of the samples. This report was presented by Academician B. P. Konstantinov. Orig. art. has: 3 figures and 1 table.

SUB CODE: 20/

SUBM DATE: 08Mar65/ ORIG REF: 008/ OTH REF: 005

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APPROVED FOR RELEASE: 08/25/2000

L 13868-66 EWT(1)/EWT(m)/EWP(t)/EWP(b) IJP(c) JD

ACC NR: AP6002425 SOURCE CODE: UR/0020/65/165/005/1062/1064

AUTHOR: Sobolev, V. V.

61

ORG: Institute of Applied Physics, Academy of Sciences MSSR (Institut prikladnoy fiziki, Akademii nauk MSSR)

TITLE: Low temperature spectral distribution of the photoconductive effect in cadmium selenide and cadmium sulfide

SOURCE: AN SSSR. Doklady, v. 165, no. 5, 1965, 1062-1064

TOPIC TAGS: cadmium selenide, cadmium sulfide, photoconductivity, single crystal, spectral distribution, low temperature effect, exciton absorption, absorption edge

ABSTRACT: Spectral curves are studied for photocurrent in CdSe and CdS at 4.2% using a monochromator with rather wide dispersion. Single crystal specimens were studied with dark currents of 10^{-9} - 10^{-11} a and photocurrents of 10^{-8} - 10^{-9} a at 77.30 and 4.20K. The spectra for the photocurrents were taken in polarized light using an instrument with a linear dispersion of 5 Å/mm and a scanning time in the 200 Å spectral region of 5-30 minutes through a slot no more than 0.5 Å wide. The data

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ACC NR: AP6002425

showed that "pure" crystals may be divided into two groups: crystals with peaks and crystals with dips on the photocurrent curves; the positions of the spectral peaks coincide with those of the dips and are highly stable. "Imperfect" crystals may also be divided into two groups. However, in this case there is a wide variation in the position and number of the peaks (or dips). The peaks (dips) in the spectral curves for the photocurrent of "pure" crystals coincide within 1-2 Å with the maxima of intense lines and bands in the line structure of the fundamental absorption edge. One of the most interesting and important results is a pronounced relationship between the structure of the photocurrent curves and the state of photoactivation of the crystals. The peaks (dips) in the photocurrent appear only when there is a certain minimum photoactivation for the specimens. There was no reliable observation of the structure of photocurrent curves in CdS at 4.2°K on a single one of the "irregular" absorption lines caused by stationary exciton-impurity complexes. The experimental data confirm the exciton origin of the peaks (diss) in the photocurrent curves for CdSe and CdS single crystals as well as the corresponding lines in the line structure of the edge absorption and emission. The exciton structure of the edge absorption shows up in the form of peaks (dips) in the spectral distribution of photocurrent because of the interaction between excitons and the photocenters which are formed when the crystal is exposed to radiation from the

Card 2/3

L 13868-66

ACC NR: AP6002425

fundamental absorption region. The results also indicate that there is no quasiequilibrium distribution in these crystals between the energy levels of different excitons (A and B) and the levels of the same exciton. Orig. art. has: 2 figures.

SUB CODE: 20/ SUBM DATE: 05Sep65/ ORIG REF: 008/ OTH REF: COO

Card 3/3 MC

. 28333-66 EWT(m)/ETC(f)/EWG(m)/T/EWP(t)/ETI LIP(c) RDW/JD ACC NR: AP6013093

SOURCE CODE: UR/0048/66/030/004/0725/0726

AUTHOR: Sobolev, V. V.

B

ORG: Institute of Applied Physics, Academy of Sciences, Moldssr (Institut priklad-

TITLE: Photoluminescence and absorption spectra of "pure" cadmium sulfide and cadmium selenide single crystals Report, Fourteenth Conference on Luminescence held in Riga 16-23 September 1965/

SOURCE: AN SSSR. Izvestiya. Seriya fizicheskaya, v. 30, no. 4, 1966, 725-726

TOPIC TAGS: photoluminescence, luminescence spectrum, absorption spectrum, cadmium selenide, cadmium sulfide

ABSTRACT: Although there have been many studies devoted to the edge absorption and luminescence spectra of CdSe and CdS, many of the details of the complex structure of these spectra are still obscure. The present work was aimed at detailed investigation of the spectra by measurements on a large number (several hundred) perfect single crystals at 4.2 and 77.3°K by means of high dispersion (3 and 6 Å/mm) spectroscopic instruments with photographic recording. The luminescence spectra of one CdSe single crystal at 4.2 and 77.3°K and of two CdS crystals at 4.2°K are reproduced on a photographic plate. The distinctive features of the absorption and emission spectra of the

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CIA-RDP86-00513R001651830004-3

L 28333-66 ACC NR: AP6013093 "pure" EdSe and CdS single crystals are described at some length and the differences are noted between the spectra of the "cold" and "warm" crystal. The wavelengths of the lines distinguished in the various spectra are listed. Information is also given on the polarization of the lines. The observed complicated structure of the exciton exciton absorption lines of CdSe and CdS can be explained on the basis of the theoretical data taking into account two molecules per unit cell and partially on the basis of the effects of spatial dispersion, the "extremum loop", and para- and ortho-excitons. Orig. art. has: 2 figures. SUB CCDE: 20/ SUEM DATE: 00/ ORIG REF: 008/ OTH REF: 006

L 31495-66 EWT(m)/ETC(f)/T/EWP(t)/ETI IJF(c) RDM/JD ACC NR: AP6013024 SOURCE CODE: UR/0051/66/020/CO4/0673/0677

AUTHOR: Sobolev, V. V.

49 B

ORG: none

TITLE: Nature of the fine structure of the absorption edge lines of single crys-

tals of cadmium selenide and sulfide

SOURCE: Optika 1 spektroskopiya, v. 20, no. 4, 1966, 673-677

TOPIC TAGS: cadmium sulfide, cadmium selenide, single crystal, absorption edge, absorption line, exciton absorption, Doppler shift, oscillation strength

ABSTRACT: The author presents a summary of his own earlier data (Opt. i spektr. v. 16, 76, 1964 and elsewhere) as well as data by others on the absorption line spectrum of single-crystal CdSe and CdS at 4.2K, obtained with very thin samples and using a spectrograph of large dispersion 3 and 6 Å/mm). The new details in the exciton absorption lines of the two substances, observed as a result, are pointed out and discussed briefly. The experimental data are compared with various theories proposed to explain this phenomenon (longitudinal excitons, extremum loops, spatial dispersion). It is concluded that a sufficiently convincing and complete explanation of the complicated structure of the exciton lines can be made

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L 31495-66

ACC NR: AP6013024

2

only by extending these theories further and by obtaining new experimental data. Among the theories proposed are also Doppler shift and splitting, realization of para-exciton states, band splitting, increase in oscillator strength, etc. The author thanks S. A. Moskalenko and M. I. Shmigly & for useful discussions and for a preprint of their paper. Orig. art. has: 2 tables.

SUB CODE: 2C/ SUBM DATE: 050ct64/ ORIG REF: 012/ OTH REF: 009

card 2/2 mc

L 46247-65 EWT(1)/T IJP(c) GG ACC NR: AP6023909

SOURCE CODE: UR/0363/66/002/007/1171/1176

AUTHOR: Sobolev, V. V.

ORG: Institute of Applied Physics, Academy of Sciences, MoldSSR (Institut prikladnoy fiziki Akademii nauk MoldSSR)

TITIE: Structure of the energy bands of certain laminar crystals

SOURCE: AN SSSR. Izv. Neorg materialy, v. 2, no. 7, 1966, 1171-1176

TOPIC TAGS: semiconductor band structure, Brillouin zone, indium compound, gallium compound, selenide, telluride

ABSTRACT: The reflection spectra of InSe, In₂Te, GaSe, and GaTe crysta⁷, were studied at T = 293 °K in the range of fundamental absorption in order to determine the character of the structure of their bands in the range E>E_g according to the band theory for laminar and tetrahedral structures, and to attempt to correlate the known optical results. The structural reflection in the range E>E_g of laminar crystals was found to be due to direct interband transitions at certain principal points of the Brillouin zone. The reflection spectra and band structure (in the range E>E_g) of GaS, GaSe, InSe, and GaTe are similar and differ markedly from those of In₂Te. The relatively simple form of the reflection spectra for E>E_g and the complex nature of the absorption spectra at E \leq E_g of the laminar crystals indicate a simple structure of their bands in the range E>E_g and a very complex structure of the lower conduction bands

Card 1/2

TMC: 548.0:531

UR/G051/66/021/001/0091/0093 EXT(m)/EWP(t)/ETI IJP(c) L 08875-67 SOURCE CODE: ACC NR: AP6025957

AUTHOR: Kroitoru, S. G.; Sobolev, V. V.

26

ORG: none

TITLE: Reflection spectra of Mg2Si and Mg2Sn crystals

SOURCE: Optika i spektroskopiya, v. 21, no. 1, 1966, 91-93

TOPIC TAGS: polycrystal, Brillouin zone, magnesium compound optic material

ABSTRACT: Reflection spectra of polished polycrystals of Mg2Si, Mg2Si, Mg2Si0.2Sn0.8, Mg₂Si_{0.5}Sn_{0.5}, Mg₂Si_{0.7}, MnSi₂, MnSi_{1.65}, and Mg₂Si_{0.6}Ge_{0.4} are studied in the range of 1 to 6 ev at 293°K. The purpose of the study was to quantitatively determine the positions of the energy zones of the crystals in various points of the Brillouin zone. All of these crystals ($Mg_2X(X=Si, Ge, Sn)$) have antifluorite lattices, are facecentered, with cubic translational symmetry. Curves of the reflection spectra are plotted and discussed in detail. Some of the peaks observed in the curves are interpreted. The authors thank Ye. N. Nikitin and N. A. Bul'onkov for supplying the crystals. Orig. art. has: 4 figures.

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CIA-RDP86-00513R001651830004-3"

ACC NRI ATGOZ4014	SOURCE CODE: UR/0000/65/000/000/0094/0104
ssledovaniya fizicheskikh (Theoretical and experiment als and other crystals). Kincopic TAGS: cadmium selent photoluminescence, temperary ABSTRACT: This is a contine sertation Abstract, Lening reflection, photoluminesce photoeffect, were observed thoroughly investigated. features of absorption and peratures 290, 160, and The with a spectrograph having sibility of observation of in the form of plates whose	the absorption of cadmium selenide single crystals out prikladnoy fiziki. Teoreticheskiye i eksperimental'nyyo svoystv poluprovodnikovykh materialov i drugikh kristallov materialov i kristope kr
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ACC NR: ATGOSHOLL

the spectra could be investigated in polarized light. Many details of the investigation and of the apparatus are described elsewhere (ZhTT v. 26, 1622, 1956; FTT v. 2, 406, 1960; DAN SSSR v. 133, 56, 1960). The experimental results and their interpretation are described in detail. The results show that the nonotonic dependence of the absorption coefficient on the wavelength of the absorbed light experiences abrupt changes (edges) in three regions of the spectrum. These edges are polarized; the edge with the longest wavelength is polarized with the electric field perpendicular to C, while the second and third edges are partially polarized. The experimental data obtained on the character of the polarization and on the magnitude of the absorption coefficient in the region of the three edges is in good agreement with the theory of the complex band structure of uniaxial crystals. The results permit identification of a number of transitions and interband distances as well as their temperature dependence. Orig. art. has: 6 figures, 3 formulas, and 3 tables.

SUB CODE: 20/ SUBM DATE: 25 Jul65/ ORIG REF: 005/ OTH REF: 004

Card 2/2

L 03008-67		
ACC NR: AP6033483	SOURCE CODE: UR/04	13/66/000/018/0087/0087
INVENTOR: <u>Semikopnyy</u> , I.		43 B
ORG: none	Av	
TITLE: Device for studyi	ng visual functions. Class 30, No.	. 186076
SOURCE: Izobret prom obr	az tov zn, no. 18, 1966, 87	:
TOPIC TAGS: visual funct	ion, vision, electric motor, human	sense, of THAL mology,
ABSTRACT: An Author Cert functions, which consists objects, and a current so and numerical indication objects, a contact mechanthe moving object, a dispostationary object, and a median object, and a current so	ificate was issued for a device, us of an electric motor, stimulus source. To provide objective examinat of results obtained in the display ism controlling frequency and durately mechanism controlling the durate chanism for setting exposure time, as of two successive stimuli (for ecce.	orces, displayed test cion of visual functions of moving and stationary cion of illumination of cion of display of the which controls duration
SUB CODE: 06/ SUBM DATE	: 08Dec64/ ATD PRESS: 5099	
Cord 1/1	UDC: 615.471.61	.2. 843.7

ACC NE. ALTERNATION

Schall College Charles and the second

AUTHOR: Andriyesh, A. M.; Sobolev, V. V.

ORG: none

TITLE: Optical reflection spectra of chalcogenides of arsenic

SOURCE: AN ESSR. Institut fiziki tverdogo tela i poluprovodnikov. Mimicheskym svyaz' v poluprovodnikakh i termodinamika (Chemical bond in semiconductors and thermodynamics). Minsk, Nauka i tekhnika, 1966, 212-216

TOPIC TAGS: arsenic compound, light reflection, glass property, optic spectrum

ABSTRACT: The purpose of the investigation was to study the change in the energy levels, in the nature and magnitude of interatomic interaction forces when compounds like As_2S_3 , As_2Se_3 , or As_2Te_3 are transformed from the crystalline state into the vitreous state. To this end the authors used spectrometers (SF-4 and SF-5) to investigate in the 1 - 6 ev region and at T=295K the reflection spectra of As_2S_3 (in the form of cleaved single crystals, glass with surface obtained by fracture, mechanically polished glass, chemically polished glass, and aged glass synthesized five years ago), and As_2Te_3 (in the form of a mechanically polished polycrystal, and glass). Plots of the reflection spectra are presented and the various irregularities observed in the spectra of the different samples are listed and interpreted. Comparison of the reflection spectra of the crystals and glasses of the different chalcogenides leads to the following conclusions: In the

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 $\mathrm{As}_2\mathrm{S}_3$ spectrum, the long-wave and short-wave maxima of the auripigment are close in position to the main peak and to the weak two maxima of artificially aged single crystal; the glass exhibits in addition to the "crystal" peaks also one structurecrystal; the glass exhibits in addition to the crystal peaks also one structureless broad band in the same region of the spectrum. In the As₂Se₃ crystal, the main tand is at 340 nm, whereas in the glass it is at 380 - 400 nm. The appearance of a peak at 340 nm in "aged" glass is due to its crystallization during the five years elapsed from the synthesis of the glass. In As2Te3, no difference is observed between the reflection of the glasses and crystals. From comparison with theoretical studies it is concluded that the maxima observed in the reflection of the singlecrystal As2S3 and As2Se3 are due to direct interband transitions. Orig. art. has: 1 figure.

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ACC NR. ATTUCTURE

COUNCE CODE: UR/OCCO/66/000/000/0221/0226

AUTHOR: Sobolev, V. V.; Syrbu, H. H.; Shutov, S. D.

ORG: none

TITLE: Energy structure of bands of certain II - V, V - VI, and III - VI compounds

SOURCE: AM BSSR. Institut fiziki tverdogo tela i poluprovodnikov. Khimicheskaya svyaz' v poluprovodnikakh i termodinamika (Chemical bond in semiconductors and thermodynamics). Minsk, Nauka i tekhnika, 1966, 221-228

TOPIC TAGS: semiconducting material, semiconductor band structure, light reflection, optic spectrum, Brillouin zone, optic transition

ABSTRACT: The purpose of the investigation was to study the reflection spectra in the region $E > E_{\rm g}$ of a large number of anisotropic semiconductors (ZnSb, CdSb, Zn₄Sb₃, Zn₃Sb₂, Cd₄Sb₃, Zn₅P₂, Cd₃P₂, Zn₅As₂, Cd₃As₂, ZnAs₂, CdAs₂, Sb₂S₃, Sb₂Se₃, Sb₂Se₃, Sb₂Te₃, Bi₂Se₃, Bi₂Te₃, InSe, In₂Te, GaSe, and GaTe) for the purpose of determining the energy gaps at different points of the Brillouin zone and comparing them with the band theories for anisotropic substances. The reflection spectra were investigated in the region 1-6 ev at T=295K. Plots of all the spectra are presented and tables of the reflection peaks for different energies are given. The main conclusion of the data is that most reflection maxima of the crystals are due to direct interband transitions; their magnitudes on the energy scale are directly equal to the values of the corresponding interband gaps at different principal points of the Brillouin zone. The

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UR/0000/66/000/000/0240/0250

AUTHOR: Kritovu, S. G.; Sobolev, V. V.; Syrbu, N. N.; Shutov, S. D.

ORG: none

TITLE: Energy band structure of crystals of groups IV, III - V, II VI, and the K32Si type

SOURCE: AN BEER. Institut fiziki tverdogo tela i poluprovodnikov. Knimicheskaya svyaz' v poluprovodnikakh i termodinamika (Chemical bond in semiconductors and thermodynamics). Minsk, Nauka i tekhnika, 1966, 240-250

TOPIC TAGS: semiconducting material, semiconductor band structure, light reflection, optic spectrum

ABSTRACT: The authors investigated the band structure, using the reflection spectra of pure and alloyed, polished and etched samples, cleaved crystals, and dendrites of groups IV and III - V, and polished and etched crystals of groups II - VI (Si, Ge, GaAs, GaSb, InAs, InSb, InP, GaP, and AlSb), Mg_Si, Mg_Si, and certain solid solutions of the systems InP-InAs, AlSb-GaSb, CdTe-HgTe, ZnSe-CdSe, Mg_Si-Mg_Si, and Mg_Si-Mg_Se. The various peaks observed on the different spectra of the substances are listed and compared with results obtained by others. Tables of the experimental values of the direct interband transitions are presented. It is stated in the conclusion that the lack of concrete and sufficiently detailed calculations of the bands and schemes for the chemical binding forces for most solids makes it very difficult

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ACC NR: AT7003885 to make further progress in the spectroscopy of crystals in k-space, which would help exp. in many physical and chemical properties of semiconductor compounds. Orig.

art. ha. 3 figures, 1 formula, and 2 tables.

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ACC NR: AT7003886

SOURCE CODE: UR/0000/66/000/000/0251/0260

Zalevskiy, B. K.; Lashkarev, G. V.; Sobolev, V. V.; Syrbu, N. N. AUTHOR:

ORG: none

TITLE: Experimental studies of the structure of energy bands in certain rare earth element chalcogenides

SOURCE: AN BSSR. Institut fiziki tverdogo tela i poluprovodnikov. Khimicheskaya svyaz' v poluprovodníkakh i termodinamika (Chemical bond in semiconductors and thermodynamics). Minsk, Nauka i tekhnika, 1966, 251-260

TOPIC TACS: compound semiconductor, refractory compound, sulfide, selenide, oxytelluride, rare earth compound, semiconductor band structure, reflection spectrum, ENERGY BAND STRUCTURE

ABSTRACT: Reflection spectra in the 200—1200 mp range of seven rare earth element chalcogenides and three oxytellurides have been obtained at 293°K and interpreted in terms of the theory of energy band structure of semiconductors. The compacted polycrystalline samples used in the experiments were prepared by sintering at 1000-1750°C powdered components in hydrogen sulfide or selenide atmosphere or in evacuated quartz ampules. Reflection spectra in the region of energy greater than the minimum forbidden energy gap (Eg) were similar for all the compounds studied. This fact indicates a great similarity in the structure of energy bands between chalcogenides and oxytellurides of the rare earth elements. Structural peculiarities

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UDC: none

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SOURCE CODE: UR/0048/6_, '030/009/1555/1557

AUTHOR: Sobolev, V. V.

ORG: Institute of Applied Physics of the Academy of Sciences of the MoldSSR (Institut prikladnoy fiziki Akademii nauk MoldSSR)

TITLE: Low-temperature photoluminescence spectra of mixed crystals of A(II)B(VI) compounds /Report, Fourteenth All-Union Conference on Luminescence (Crystal Phosphors) held at Riga, 16-23 Sept. 1965/

SOURCE: AN SSSR. Izvestiya. Seriya fizicheskaya, v.30, no.9, 1966, 1555-1557

TOPIC TAGS: luminescence spectrum, band spectrum, line spectrum, zinc compound, cadmium compound sulfide, selenide, telluride, mixed crystal

ABSTRACT: The author has recorded the photoluminescence spectra at 4.2° K of CdS.CdSe single crystals of two different compositions and of coarse-grained mixed polycrystalline layers of each of the 10 systems containing two of the following compounds: ZnS, ZnTe, CdS, CdTe, and CdSe. The polycrystalline systems were investigated over a wide range of compositions. The principal luminescence bands are tabugated over a wide range of compositions. The principal luminescence bands are tabugated. All the investigated systems except those containing CdTe exhibited banded lated. All the equidistant bands. The failure of the CdTe-containing systems

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of the M_2X_3 and MX compound semiconductors were derived from the weak reflection peaks of Ce_2S1_3 , Nd_2S1_3 , and EuSe and from the reflection peaks in the 240—420 mu region of Sm_2S_3 and sesquiselenides of Le, Le, PR, Nd, and Sm. Orig. art. has: 4 figures, 1 table, and 3 formulas.

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lattice constants of CdTe and those of the other materials. In some of the systems the energy separation of the equidistant edge luminescence bands was equal to the energy of a longitudinal phonon, and in others of them it was not. Line spectra of edge luminescence were observed in those of the polycrystalline CdSe.CdS layers that did not contain more than about 33% of CdS, but line spectra were not observed in the CdSe.CdS single crystals. The number and sharpness of the edge luminescence lines decreased with increasing CdS content. The strongest edge luminescence lines are ascribed to resonance Ausleuchtung from the lowest exciton states, and the other lines are ascribed mainly to excitons bound in the mixed crystal lattice. The absence of lines in the edge luminescence spectra of the single crystals is ascribed to the presence in the crystals of many centers at which there take place nonradiative exciton and electron-hole processes. Orig. art. has: 2 figures and 1 table. OTH REF: 001

ORIG. REF: 007 none SUBM DATE: SUB CODE:

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ACC NRI AP7005003 (A) SOURCE CODE: UR/0048/66/03	0/009/1558/1559
AUTHOR: Sobolev, V.V. ORG: Institute of Applied Physics of the Academy of Sciences of the Academy of Sc	f the MoldSSR

(Institut prikladnoy fiziki Akademii nauk MoldSSR)

TITLE: Effects of impurities, heat treatment, and deformation on the luminescence and absorption spectra of cadmium sulfide and cadmium selenide /Report, Fourteenth All-Union Conference on Luminescence (Crystal Phosphors) held at Riga, 16-23 Sept. 1965/

SOURCE: AN SSSR. Izwestiye, Seriya fizicheskaya, v.30, no.9, 1966, 1558-1559

TOPIC TAGS: cadmium sulfide, cadmium selenide, luminescence spectrum, absorption spectrum, mechanical stress, heat effect, crystal impurity, single crystal

ABSTRACT: The author investigated the effects of different impurities, heat treatment at 700-800° C in vacuum and in different atmospheres, and mechanical stress on the edge luminescence, absorption, and reflection spectra of CdSe and CdS single crystals at 4.20 K. In the present paper the results are discussed qualitatively and two of the spectra are presented as examples. The heat treatment resulted in a broadening of the absorption spectrum, as though there had arisen an additional continuous absorption, and some smearing and shift of the line structure and the absorption edge. Different impurities and different effects on the absorption spectra, and some of them (Br and Cl) had no appreciable effect on the CdSe absorption spectrum

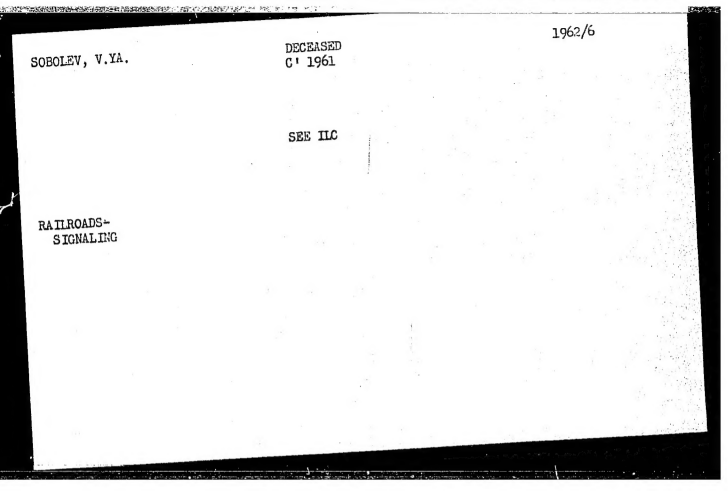
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Mechanical stress was introduced by cementing the specimens to different substrates and subsequently cooling them. The effect of strain on the absorption spectra was very similar to that of heat treatment, and it is concluded that the effect of heat treatment is due to the introduction of internal deformations rather than to changes in the absorption at impurities and defects. Both CdSe and CdS are very sensitive to lattice distortions, and this sensitivity may find technical application. Heating the crystals, particularly in an atmosphere of the anion vapor, considerably altered the edge luminescence spectra, distorting the continuous edge luminescence spectrum and reducing the intensity of the edge luminoscence lines. The presence of impurities greatly altered the luminescence spectra, resulting in the appearance of many narrow and sometimes strong lines between 6810 and 7100 A and in enhancement of the doublet structure of the edge luminescence bands. On the basis of the present and other results it is concluded that "pure" CdSe crystals grown by the Frerichs technique are much more nearly pure than are the corresponding "pure" CdS crystals. Many details of the edge luminescence spectra can be explained on the basis of the exciton and exciton - impurity center models, but they are not so explained in the present paper. Orig. art. has: 2 figures.

SUB CODE: 20 SUBM DATE: none ORIG. RKF: 003

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